What is claimed is:

- 1. A method for analyzing a complex biological sample using a Fourier Transform Mass Spectrometer (FTMS), said method comprising the steps of:
 - a. ionizing a sample to produce sample (molecular) ions;
 - b. introducing said ions into an analysis region of said FTMS;
 - c. analyzing said ions to determine the molecular weight and abundance of said ions;
 - d. utilizing said molecular weight to determine the empirical formula of each species of said sample; and
 - e. identifying each said species by comparing said empirical formula to a database of formulas for known molecules.
- 2. A method according to claim 1, wherein said determining of the molecular weight is performed with an accuracy sufficient to identify empirical formula of said ions.
- 3. A method according to claim 1, wherein said database of known molecules is updated with said determined molecular structures.

- 4. A method for analyzing a complex biological sample using a Fourier Transform Mass Spectrometer (FTMS), said method comprising the steps of:
 - a. ionizing a sample to produce sample (molecular) ions;
 - b. introducing said ions into an analysis region of saidFTMS;
 - c. analyzing said ions to determine the molecular weight and abundance of said ions;
 - d. determining the molecular structure of each species by multiple stages of mass spectrometry; and
 - e. producing a profile of the sample showing structure and concentration data for each species.
- 5. A method according to claim 4, wherein said determining of the molecular weight is performed with an accuracy sufficient to identify empirical formula of said ions.

- 6. A method for analyzing a complex biological sample using a Fourier Transform Mass Spectrometer (FTMS), said method comprising the steps of:
 - a. ionizing the sample to produce sample precursor ions;
 - b. introducing said ions into the analysis region of said FTMS;
 - c. analyzing said ions to determine the molecular weight, the abundance and the empirical formula of said ions;
 - d. fragmenting said sample precursor ions to produce fragment ions;
 - e. determining the molecular weight, the abundance and empirical formula of said fragment ions;
 - f. determining the structure of said fragment ions by comparing said empirical formulas of said fragment ions to a database of fragments with known structure;
 - g. combining said structures of said fragment ions to determine the precursor ion structure for each species in said sample; and
 - h. producing a profile of said sample showing structure and concentration data for selected species of said sample.

- 7. A method according to claim 6, wherein said determining of the molecular weight is performed with an accuracy sufficient to identify empirical formula of said ions.
- 8. A method according to claim 6, wherein said fragmenting is performed using photodissociation.
 - 9. A method for analyzing a complex biological sample utilizing
 Fourier Transform Mass Spectrometry (FTMS), said method
 comprising the steps of:
 - a. ionizing polar molecules using positive and negative electrospray to produce sample (molecular) ions;
 - b. introducing said ions into an analysis region of said FTMS;
 - c. analyzing said ions to determine the molecular weight and abundance of said ions;
 - d. utilizing said molecular weight to determine the empirical formula of each species of said sample; and
 - e. identifying each said species by comparing said empirical formula to a database of formulas for known molecules.

- 10. A method according to claim 9, wherein said determining of the molecular weight is performed with an accuracy sufficient to identify empirical formula of said ions.
- 11. A method according to claim 9, wherein said database of known molecules is updated with said determined molecular structures.
 - 12. A method for analyzing a complex biological sample utilizing Fourier Transform Mass Spectrometry (FTMS), said method comprising the steps of:
 - a. ionizing non-polar molecules using positive and negative ion atmospheric pressure chemical ionization to produce sample (molecular) ions.
 - b. introducing said ions into an analysis region of saidFTMS;
 - c. analyzing said ions to determine the molecular weight and abundance of said ions;
 - d. utilizing said molecular weight to determine the empirical formula of each species of said sample; and
 - e. identifying each said species by comparing said empirical formula to a database of formulas for known molecules.

- 13. A method according to claim 12, wherein said determining of the molecular weight is performed with an accuracy sufficient to identify empirical formula of said ions.
- 14. A method according to claim 12, wherein said database of known molecules is updated with said determined molecular structures.